

FORM PTO-1390 (REV 10-2000)		U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE		ATTORNEY'S DOCKET NUMBER
<b>TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371</b>		CU-2652 RJS		
		U.S. APPLICATION NO. (If known, see 37 CFR 1.5) <b>09/936746</b>		
INTERNATIONAL APPLICATION NO. PCT/EP00/01837	INTERNATIONAL FILING DATE 03 March 2000	PRIORITY DATE CLAIMED 12 March 1999		
TITLE OF INVENTION COSMETIC PREPARATIONS				
APPLICANT(S) FOR DO/EO/US Ute GRIESBACH et al				
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:				
<ol style="list-style-type: none"> <li><input checked="" type="checkbox"/> This is a <b>FIRST</b> submission of items concerning a filing under 35 U.S.C. 371.</li> <li><input type="checkbox"/> This is a <b>SECOND</b> or <b>SUBSEQUENT</b> submission of items concerning a filing under 35 U.S.C. 371.</li> <li><input checked="" type="checkbox"/> This is an express request to promptly begin national examination procedures (35 U.S.C. 371(f)).</li> <li><input checked="" type="checkbox"/> The US has been elected by the expiration of 19 months from the priority date (PCT Article 31).</li> <li><input checked="" type="checkbox"/> A copy of the International Application as filed (35 U.S.C. 371(c)(2))           <ol style="list-style-type: none"> <li><input type="checkbox"/> is attached hereto (required only if not communicated by the International Bureau).</li> <li><input checked="" type="checkbox"/> has been communicated by the International Bureau.</li> <li><input type="checkbox"/> is not required, as the application was filed in the United States Receiving Office (RO/US).</li> </ol> </li> <li><input type="checkbox"/> An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)).</li> <li><input type="checkbox"/> Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3))           <ol style="list-style-type: none"> <li><input type="checkbox"/> are attached hereto (required only if not communicated by the International Bureau).</li> <li><input type="checkbox"/> have been communicated by the International Bureau.</li> <li><input type="checkbox"/> have not been made; however, the time limit for making such amendments has NOT expired.</li> <li><input type="checkbox"/> have not been made and will not be made.</li> </ol> </li> <li><input type="checkbox"/> An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).</li> <li><input type="checkbox"/> An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).</li> <li><input type="checkbox"/> An English language translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).</li> </ol>				
Items 11 to 16 below concern document(s) or information included:				
<ol style="list-style-type: none"> <li><input type="checkbox"/> An Information Disclosure Statement under 37 CFR 1.97 and 1.98.</li> <li><input type="checkbox"/> An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.</li> <li><input type="checkbox"/> A <b>FIRST</b> preliminary amendment.           <input type="checkbox"/> A <b>SECOND</b> or <b>SUBSEQUENT</b> preliminary amendment.</li> <li><input type="checkbox"/> A substitute specification.</li> <li><input type="checkbox"/> A change of power of attorney and/or address letter.</li> <li><input type="checkbox"/> Other items or information:</li> </ol>				
Express Mail Label No.: EL698182332US				

17.  The following fees are submitted:

## BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5) ) :

Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO ..... \$1000.00

International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO ..... \$860.00

International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO ..... \$710.00

International preliminary examination fee paid to USPTO (37 CFR 1.482) but all claims did not satisfy provisions of PCT Article 33(1)-(4) ..... \$690.00

International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(1)-(4) ..... \$100.00

ENTER APPROPRIATE BASIC FEE AMOUNT =

\$ 860.00

Surcharge of \$130.00 for furnishing the oath or declaration later than  20  30 months from the earliest claimed priority date (37 CFR 1.492(e)).

\$

CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE	
Total claims	- 20 =		X \$18.00	\$
Independent claims	- 3 =		X \$80.00	\$
MULTIPLE DEPENDENT CLAIM(S) (if applicable)			+ \$270.00	\$
TOTAL OF ABOVE CALCULATIONS =				\$ 860.00
<input checked="" type="checkbox"/> Applicant claims small entity status. See 37 CFR 1.27. The fees indicated above are reduced by 1/2.				\$ 430.00
SUBTOTAL =				\$
Processing fee of \$130.00 for furnishing the English translation later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(f)).				\$
+				
TOTAL NATIONAL FEE =				\$
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property				\$
+				
TOTAL FEES ENCLOSED =				\$ 430.00
				Amount to be refunded: \$
				charged: \$

a.  A check in the amount of \$ 430.00 to cover the above fees is enclosed.

b.  Please charge my Deposit Account No. 12-0400 in the amount of \$ 430.00 to cover the above fees. A duplicate copy of this sheet is enclosed.

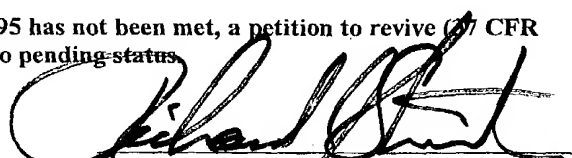
c.  The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 12-0400. A duplicate copy of this sheet is enclosed.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

SEND ALL CORRESPONDENCE TO:

Ladas & Parry  
224 South Michigan Avenue  
Suite 1200  
Chicago, Illinois 60604  
(312) 427-1300

September 12, 2001



SIGNATURE:  
Richard J. Streit

NAME  
25765

REGISTRATION NUMBER

DOCKET: CU-2652

***IN THE UNITED STATES PATENT & TRADEMARK OFFICE***

APPLICANT: Ute GRIESBACH et al )  
S SERIAL NO: 09/936,746 )  
TITLE: COSMETIC PREPARATIONS )  
COMPLETION OF PCT/EP00/01837 filed 03 March 2000 )

The Commissioner for Patents (DO/EO/US)  
Box PCT  
Washington, D.C. 20231

**PRELIMINARY AMENDMENT**

Dear Sir:

This is a preliminary amendment which corrects minor deficiencies in the claims as filed.

**IN THE CLAIMS:**

Please replace claims 2-9 with the attached clean version of replacement claims 2-9. Please see a marked up version of the amendment and claims attached hereto to aid the Examiner in identification of the changes.

**REMARKS**

Applicants are submitting the claims to better clarify them for prosecution in the United States.

If the Examiner has any questions, the Examiner may contact the undersigned at the listed telephone number.

Respectfully submitted,

March 5, 2002

Date

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Ute Griesbach et al  
U.S. Serial No. 09/936,746  
Docket: CU-2652  
Clean Version of Amended Claims

We claim

1. Cosmetic preparations, containing
  - (a) water soluble  $\beta$ -(1,3) glucans, substantially free from  $\beta$ -(1,6) linkages, and
  - (b) chitosans.
2. Preparations according to claim 1, which contain glucans which are obtained based on yeasts from the family *Saccharomyces*.
3. Preparations according to claim 1, which contain glucans which are obtained by contacting glucans with  $\beta$ -(1,3) and  $\beta$ -(1,6) linkages with  $\beta$ -(1,6) glucanases, in such a way that practically all  $\beta$ -(1,6) linkages are loosened.
4. Preparations according to claim 3, wherein the glucans which are used, previously have been treated with glucanases based on *Trichoderma harzianum*.
5. Preparations according to claim 1, which contain chitosans with molecular weights in the area from 50 000 to 500 000 Daltons.
6. Preparations according to claim 1, which contain chitosans with molecular weights in the area from 800 000 to 1 200 000 Daltons.
7. Preparations according to claim 1, which contain carboxylated chitosans.
8. Preparations according to claim 1, which contain succinylated chitosans.

## 9. Preparations according to claim 1, which contain

- (a) 0.01 to 25 % by weight of water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages, and
- (b) 0.01 to 5 % by weight of chitosans,

provided that the stated amounts are supplemented with water as well as optionally other auxiliaries and additional agents up 100 % by weight.

## 10. Use of mixtures containing

- (a) water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages,
- (b) chitosans,

for manufacturing of cosmetic preparations.

Ute Griesbach et al  
U.S. Serial No. 09/936,746  
Docket: CU-2652  
Marked Version of Amended Claims

5

**[P a t e n t c l a i m s] We claim:**

1. Cosmetic preparations, containing
  - 10 (a) water soluble  $\beta$ -(1,3) glucans, substantially free from  $\beta$ -(1,6) linkages, and
  - (b) chitosans.
2. Preparations according to claim 1, **[characterised by that they] which** contain glucans which are obtained based on yeasts from the family
  - 15 *Saccharomyces*.
3. Preparations according to **[claim 1 and/or 2, characterised by that they]** **claim 1, which** contain glucans which are obtained by contacting glucans with  $\beta$ -(1,3) and  $\beta$ -(1,6) linkages with  $\beta$ -(1,6) glucanases, in such a way that practically
  - 20 all  $\beta$ -(1,6) linkages are loosened.
4. Preparations according to claim 3, **[characterised by that] wherein the** glucans **which** are used, **[which]** previously have been treated with glucanases based on *Trichoderma harzianum*.
  - 25
5. Preparations according to **[at least one of the claims 1 to 4, characterised by that they]** **claim 1, which** contain chitosans with molecular weights in the area from 50 000 to 500 000 Daltons.
- 30 6. Preparations according to **[at least one of the claims 1 to 4, characterised by that they]** **claim 1, which** contain chitosans with molecular weights in the area from 800 000 to 1 200 000 Daltons.

7. Preparations according to [at least one of the claims 1 to 6, characterised by that they] claim 1, which contain carboxylated chitosans.

8. Preparations according to [at least one of the claims 1 to 7, 5 characterised by that they] claim 1, which contain succinylated chitosans.

9. Preparations according to [at least one of the claims 1 to 8, characterised by that they] claim 1, which contain  
10 (a) 0.01 to 25 % by weight of water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages, and  
(b) 0.01 to 5 % by weight of chitosans,  
provided that the stated amounts are supplemented with water as well as optionally other auxiliaries and additional agents up 100 % by weight.

15 10. Use of mixtures containing  
(a) water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages,  
(b) chitosans,  
for manufacturing of cosmetic preparations.

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## **PATENT APPLICATION**

PCT/EP00/01837 (00.03.03)

**APPLICANT:** BIOTEC ASA  
N-STRANDGT. 3  
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NORWAY

## **TITLE: "Cosmetic preparations"**

## COSMETIC PREPARATIONS

### The field of the invention

The invention belongs to the field of cosmetics and concerns preparations, especially for the treatment of the skin and hair, which contain a synergistic mixture of specific water soluble  $\beta$ -glucans and chitosans, as well as the use of the mixtures for the production of cosmetic materials.

### Prior art

10 The formation of wrinkles caused by increasing age is induced through the degradation of different macro molecules such as for example elastin and collagen, which are responsible for the elastases. Many inflammatory skin diseases, such as for example psoriasis or UV erythema, can also be causatively 15 linked to an increased concentration of serine proteases, such as e.g. elastase in the upper skin areas [see R.Voegeli et al. in *Cosm. Toil.* 111, 51(1996)].

The formation of wrinkles in the skin is normally not counteracted by means of physiological active principles, but by means of cosmetic agents. Many so-called "anti-ageing products" contain liposomes loaded with water or aqueous 20 active agents, which through the fat layer of the skin are reaching the epidermis, where they gradually dissolve and through continuous water release compensate the skin recesses and regulate the moisture content of the skin. However, this effect is no combat against the causes, but only has a so-called "repair effect", which lasts only for a short period of time. Also the use of specific 25 polysaccharides as agents against the skin ageing is known from prior art. Thus it has been suggested in the patent US 5,223,491 to employ a carboxymethylated  $\beta$ -1,3 glucan, which had been extracted from the yeast fungus *Saccharomyces cerevisiae*, for topical application. The glucan is, however, insoluble in water and can accordingly only be formulated with much difficulties. From the European 30 patent EP-B1 0500718 (Donzis) is further the use of water insoluble  $\beta$ -(1,3) glucans, which are obtained from the cell walls of yeast, known for revitalization of the skin. Finally, in WO 98/40082 (Henkel) the use of water soluble  $\beta$ -(1,3) glucans as active agents for the treatment of the skin have been proposed. But also these glucans, which preferably are schizophyllan or krestin, i.e. extracts of 35 fungi, have in practice not shown to be sufficiently effective.

The task of the present invention was therefore to make available novel cosmetic agents, which distinguish themselves in the field of skin treatment through an improved vitalization. Especially skin ageing, formation of wrinkles and skin roughness should be improved.

5

### Description of the Invention

The object of the invention are cosmetic preparations, containing

10 (a) water soluble  $\beta$ -(1,3) glucans, substantially free from  $\beta$ -(1,6) linkages, and  
(b) chitosans.

Surprisingly it was found, that the addition of chitosans increases the skin vitalizing properties of specific  $\beta$ -(1,3) glucans in a synergistic way, while conversely the specific  $\beta$ -(1,3) glucans definitive improve the film forming 15 properties of chitosans. In this manner it is possible especially to produce agents for skin and hair treatments, but also agents for sun protection with special performance properties.

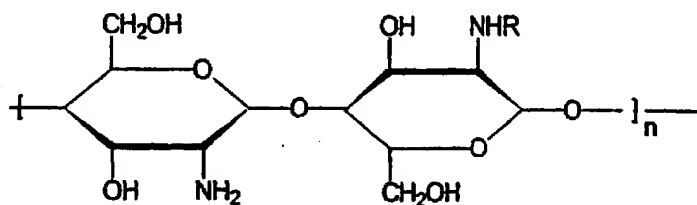
### Water soluble $\beta$ -(1,3) glucans

20 The term glucans is intended to mean homopolysaccharides based on glucose. Depending on sterical linking there is a difference between  $\beta$ -(1,3),  $\beta$ -(1,4) and  $\beta$ -(1,6) glucans.  $\beta$ -(1,3) Glucans normally show a helical structure, whereas glucans with a (1,4) linkage generally have a linear structure. The  $\beta$ -glucans of the invention have a (1,3) structure, i.e. they are substantially free 25 from undesired (1,6) linkages. Preferably such  $\beta$ -(1,3) glucans are used where the side chains exclusively show (1,3) linkages. Especially the agents contain glucans which are obtained on the basis of yeast from the family *Saccharomyces*, especially *Saccharomyces cerevisiae*. Glucans of this type are available in technical amounts according to known methods. The international patent 30 application WO 95/30022 (Biotec-Mackzymal) describes e.g. a method for producing such substances, wherein glucans with  $\beta$ -(1,3) and  $\beta$ -(1,6) linkages are brought in contact with  $\beta$ -(1,6) glucanases in such a way, that practically all  $\beta$ -(1,6) linkages are loosened. Preferably used for the manufacture of these glucans are glucanases based on *Trichoderma harzianum*. As to the manufacture

and availability of the glucans contained in these agents, reference is made to the above cited publication.

### Chitosans

5 Chitosans are biopolymers and belong to the group of hydrocolloids. From a chemical point of view they are partial deacetylated chitins with different molecular weights, and contain the following - idealized - monomer module:



In contrast to most of the hydrocolloids, which are negatively charged in the 10 range of biological pH-values, chitosans are under these conditions cationic biopolymers. The positively charged chitosans can interact with opposite charged surfaces and are therefore used in cosmetic hair and body care agents as well as in pharmaceutical preparations (see *Ullmann's Encyclopedia of Industrial Chemistry*, 5th Ed., vol. A6, Weinheim, Verlag Chemie, 1986, p. 231-332). A 15 summary of these subjects are also published in for example B. Gesslein et al., *HAPPI* 27, 57 (1990), O. Skaugrud in *Drug Cosm. Ind.* 148, 24 (1991) and E. Onsoyen et al. in *Seifen-Öle-Fette-Wachse* 117, 633 (1991). By the production of chitosan chitin is used as starting material, preferably the shell residues of crust animals, which are available in large amounts as cheap raw materials. The chitin 20 is thereby, using a method which first was described by Hackmann et al., usually first deprotonated by addition of bases, demineralized by addition of mineral acids and at last deacetylated by addition of strong bases, whereby the molecular weights can be distributed over a broad spectrum. Corresponding methods are for example known from *Makromol. Chem.* 177, 3589 (1976) or the French patent 25 application FR-A1 2701266. Preferably use is made of such types which are described in the German patent applications DE-A1 4442987 and DE-A1 19537001 (Henkel), and which have an average molecular weight of 10 000 to 2 500 000, preferably 800 000 to 1 200 000 Daltons, a viscosity according to Brookfield (1 % by weight in glycolic acid) below 5 000 mPas, a degree of

deacetylation in the range of 80 to 88 % and a content of ashes of less than 0,3 % by weight. In addition to the chitosans as typical cationic biopolymers come according to the invention also in question anionic, respectively nonionic derivatized chitosans, such as e.g. carboxylation, succinilation or alkoxylation products, as they are described for example in the German patent DE-C2 5 3713099 (L'Oreal) as well as in the German patent application DE-A1 19604180 (Henkel).

In a preferable embodiment of the invention, the preparations contain  
(a) 0,01 to 25, preferably 0,5 to 20 and especially 1 to 5 % by weight of water  
10 soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages, and  
(b) 0,01 to 5, preferably 0,5 to 3 and especially 1 to 2 % by weight of chitosans,  
provided that the used amounts together with water and possibly other auxiliary  
15 and additional substances summarize to 100 % by weight.

#### Commercial applicability

The addition of chitosans leads to a synergistic increase in the skin  
20 vitalizing properties and film forming properties of glucans. A further object of the present invention concerns the use of mixtures which contain  
(a) water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages, and  
(b) chitosans,  
25 for production of cosmetic preparations, especially care and cleaning agents for skin and hair, as well as sun protection agents.

The preparations according to the invention, such as e.g. hair shampoos, hair lotions, foam baths, sun protection agents, lotions or cremes for face and body care, baby care products, decorative cosmetics, gels or ointments and  
30 suchlike can further as additional auxiliary or additional substances contain mild surfactants, oil bodies, emulsifiers, hyperfattening agents, pearl lustre waxes, consistency substances, thickening agents, polymers, silicon compounds, fats, waxes, stabilizing agents, biogenic active substances, deodorants, agents against dandruff, film forming agents, swelling agents, UV light protection factors,  
35 antioxidants, inorganic colour pigments, hydrotropes, preservatives, insect

repellents, self tanning agents, solubilizing agents, perfume oils, colouring agents and suchlike.

Typical examples of suitable mild, i.e. especially skin compatible **surfactants**, are fatty alcohol polyglycol ether sulphates, monoglyceride sulphates, mono- and/or dialkyl sulfosuccinates, fatty acid isethionates, fatty acid sarcosinates, fatty acid taurides, fatty acid glutamates,  $\alpha$ -olefine sulphonates, ethercarboxylic acids, alkyl oligoglucosides, fatty acid glucamides, alkylamido betaines and/or protein fatty acid condensates, the last mentioned preferably based on wheat proteins.

As **oil bodies** use can be made of for example Guerbet alcohols based on fatty alcohols with 6 to 18, preferably 8 to 10 carbon atoms, esters of linear C<sub>6</sub>-C<sub>22</sub> fatty acids with linear C<sub>6</sub>-C<sub>22</sub> fatty alcohols, esters of branched C<sub>6</sub>-C<sub>13</sub> carboxylic acids with linear C<sub>6</sub>-C<sub>22</sub> fatty alcohols, such as e.g. myristyl myristate, myristyl palmitate, myristyl stearate, myristyl isostearate, myristyl oleate, myristyl behenate, myristyl erucate, cetyl myristate, cetyl palmitate, cetyl stearate, cetyl isostearate, cetyl oleate, cetyl behenate, cetyl erucate, stearyl myristate, stearyl palmitate, stearyl stearate, stearyl isostearate, stearyl oleate, stearyl behenate, stearyl erucate, isostearyl myristate, isostearyl palmitate, isostearyl stearate, isostearyl isostearate, isostearyl oleate, isostearyl behenate, isostearyl oleate, oleyl myristate, oleyl palmitate, oleyl stearate, oleyl isostearate, oleyl oleate, oleyl behenate, oleyl erucate, behenyl myristate, behenyl palmitate, behenyl stearate, behenyl isostearate, behenyl oleate, behenyl behenate, behenyl erucate, erucyl myristate, erucyl palmitate, erucyl stearate, erucyl isostearate, erucyl oleate, erucyl behenate and erucyl erucate. In addition esters of linear C<sub>6</sub>-C<sub>22</sub> fatty acids with branched alcohols, especially 2-ethylhexanol, esters of hydroxycarboxylic acids with linear or branched C<sub>6</sub>-C<sub>22</sub> fatty alcohols, especially dioctyl malate, esters of linear and/or branched fatty acids with polyvalent alcohols (such as e.g. propylene glycol, dimeric diol or trimeric triol) and/or Guerbet alcohols, triglycerides based on C<sub>6</sub>-C<sub>10</sub> fatty acids, liquid mixtures of mono-/di-/triglycerides based on C<sub>6</sub>-C<sub>18</sub> fatty acids, esters of C<sub>6</sub>-C<sub>22</sub> fatty alcohols and/or Guerbet alcohols with aromatic carboxylic acids, especially benzoic acid, esters of C<sub>2</sub>-C<sub>12</sub> dicarboxylic acids with linear or branched alcohols with 1 to 22 carbon atoms or polyols with 2 to 10 carbon atoms and 2 to 6 hydroxyl groups, plant oils, branched primary alcohols, substituted cyclohexanes, linear and branched C<sub>6</sub>-C<sub>22</sub> fatty

alcohol carbonates, Guerbet carbonates, esters of benzoic acid with linear and/or branched C<sub>6</sub>-C<sub>22</sub> alcohols (e.g. Finsolv® TN), linear or branched, symmetrical or unsymmetrical dialkyl ethers with 6 to 22 carbon atoms in each alkyl group, ring opening products of epoxydized fatty acid esters with polyols, silicone oils and/or 5 aliphatic or naphthenic hydrocarbons, such as e.g. squalan, squalen or dialkyl cyclohexanes, can be used

As **emulsifiers** for example nonionic surfactants from at least one of the following groups may be used:

- (1) Addition products of 2 to 30 moles ethylene oxide and/or 0 to 5 moles 10 propylene oxide on linear fatty alcohols with 8 to 22 C atoms, on fatty acids with 12 to 22 C atoms and on alkyl phenols with 8 to 15 C atoms in the alkyl group;
- (2) C<sub>12/18</sub> fatty acid mono- and -diesters of addition products of 1 to 30 moles ethylene oxide and glycerol;
- 15 (3) glycerol mono- and diesters and sorbitan mono- and diesters of saturated and unsaturated fatty acids with 6 to 22 carbon atoms and their ethylene oxide addition products;
- (4) alkyl mono- and oligoglycosides with 8 to 22 carbon atoms in the alkyl group and their ethoxylated analogues;
- 20 (5) addition products of 15 to 60 moles ethylene oxide on ricinus oil and/or hardened ricinus oil;
- (6) polyol and especially polyglycerol esters, such as e.g. polyglycerol polyricinoleate, polyglycerol poly-12-hydroxystearate or polyglycerol dimerate isostearate, and also mixtures of compounds from more of these classes of 25 substances;
- (7) addition products of 2 to 15 moles ethylene oxide on ricinus oil and/or hardened ricinus oil;
- (8) partial esters based on linear, branched, unsaturated or saturated C<sub>6/22</sub> fatty acids, ricinolic acid and 12-hydroxy stearic acid and glycerol, polyglycerol, 30 pentaerythrone, dipentaerythrone, sugar alcohols (e.g. sorbitol), alkyl glucosides (e.g. methyl glucoside, butyl glucoside, lauryl glucoside) as well as polyglucosides (e.g. cellulose);
- (9) mono-, di- and trialkylphosphates as well as mono-, di- and/or tri-PEG alkylphosphates and their salts;

- (10) wool wax alcohols;
- (11) polysiloxane/polyalkyl/polyether copolymers or corresponding derivatives;
- (12) mixed esters of pentaerythrite, fatty acids, citric acid and fatty alcohol according to DE 1165574 PS and/or mixed esters of fatty acids with 6 to 22 carbon atoms, methyl glucose and polyols, preferably glycerol or polyglycerol,
- (13) polyalkylene glycols, as well as
- (14) glycerol carbonate.

The addition products of ethylene oxide and/or of propylene oxide on fatty alcohols, fatty acids, alkyl phenols, glycerol mono- and diesters as well as sorbitan mono- and -diesters of fatty acids or on ricinus oil are known products which are commercially available. They are mixtures of homologous substances, with average degree of alkoxylation corresponding to the ratio of the amounts of the substances ethylene oxide and/or propylene oxide and substrate, with which the addition reaction is carried out. C<sub>12/18</sub> fatty acid mono- and -diesters of addition products of ethylene oxide on glycerol are known from DE 2024051 PS as revertive fatting agents for cosmetic preparations.

C<sub>8/18</sub> alkyl mono- and oligoglycosides, their manufacture and their use is known from prior art. Their preparation can especially be carried out by reaction of glucose or oligosaccharides with primary alcohols having 8 to 18 C atoms. With regard to the glycoside residue both monoglycosides, where a cyclic sugar group is glycosidic bond to the fatty alcohol, and oligomeric glycosides with a degree of oligomerisation until preferably about 8, are suitable. The degree of oligomerization is then a statistical mean value, based on a distribution of homologues which is usual for such products of technical quality.

Zwitterionic surfactants can also be used as emulsifiers. The term zwitterionic surfactants is intended to mean such surface active compounds which in their molecule have at least a quaternary ammonium group and at least one carboxylate and one sulphonate group. Especially suitable zwitterionic surfactants are the so-called betaines such as the N-alkyl-N,N-dimethyl ammonium glycinate, for example the coco alkyltrimethyl ammonium glycinate, N-acylaminopropyl-N,N-dimethyl ammonium glycinate, for example the coco acylaminopropyl dimethyl ammonium glycinate, and 2-alkyl-3-carboxymethyl-hydroxyethyl imidazoline with in each case 8 to 18 C atoms in the alkyl or acyl -

groups, as well as the coco acylaminoethyl hydroxyethylcarboxymethyl glycinate. Especially preferred is that under the CTFA term *cocamidopropyl betaine* known fatty acid amide derivative. Also suitable emulsifiers are ampholytic surfactants. Ampholytic surfactants are such surface active compounds which in addition to a 5  $C_{8/18}$  alkyl or acyl group in the molecule at least contain a free amino group and at least one -COOH or -SO<sub>3</sub>H group and which can form inner salts. Examples of suitable ampholytic surfactants are N-alkyl glycines, N-alkyl propionic acids, N-alkyl aminobutyric acids, N-alkyl iminodipropionic acids, N-hydroxyethyl-N-alkylamidopropyl glycines, N-alkyltaurines, N-alkylsarcosines, 10 2-alkylaminopropionic acids and alkylamino acetic acids with in each case about 8 to 18 C atoms in the alkyl group. Especially preferable ampholytic surfactants are the N-coco alkylamino propionate, the coco acylamino ethylaminopropionate and the  $C_{12/18}$  acylsarcosine. In addition to the ampholytic, also quaternary emulsifiers can be used, of which ester salts of the type of esterquats, preferably 15 methylquaternised di-fatty acid triethanolamine ester salts, are especially preferable.

As **hyperfattening** agents substances such as for example lanolin and lecithin as well as polyethoxylated or acylated lanolin and lecithin derivatives, polyol fatty acid esters, monoglycerides and fatty acid alkanolamides can be used, 20 whereby the last mentioned at the same time act as foam stabilisers.

As exemplary **pearl gloss waxes** the following should be mentioned: Alkylene glycolester, especially ethyleneglycol distearate; fatty acid alkanolamides, especially coco fatty acid diethanolamide; partial glycerides, especially stearic acid monoglyceride; esters of polyvalent, possibly 25 hydroxysubstituted carboxylic acids with fatty alcohols with 6 to 22 carbon atoms, especially long chain esters of tartaric acid; fat substances, such as for example fatty alcohols, fatty ketones, fatty aldehydes, fatty ethers and fatty carbonates, wherin the sum of carbon atoms is at least 24, especially lauron and distearylether; fatty acids such as stearic acid, hydroxystearic acid or behenic 30 acid, ring opening products of olefine epoxides with 12 to 22 carbon atoms with fatty alcohols with 12 to 22 carbon atoms and/or polyols with 2 to 15 carbon atoms and 2 to 10 hydroxyl groups as well as their mixtures.

As **consistency givers** preferably use is made of fatty alcohols or hydroxy fatty alcohols with 12 to 22 and preferably 16 to 18 carbon atoms and additionally

partial glycerides, fatty acids or hydroxy fatty acids. A combination of these substances with alkyl oligoglucosides and/or fatty acid-N-methyl glucamides with the same chain length and/or polyglycerol-poly-12-hydroxy stearates.

Suitable **thickening agents** are for example types of aerosil (hydrophilic 5 silicic acids), polysaccharides, especially xanthan gum, guar-guar, agar-agar, alginates and tyloses, carboxymethyl celluloses and hydroxyethyl celluloses, as well as higher molecular polyethylenglycol mono- and diesters of fatty acids, polyacrylates, (e.g. Carbopol® from Goodrich or Synthalenes® from Sigma), polyacrylamides, polyvinyl alcohol and polyvinyl pyrrolidone, surfactants such as for 10 example ethoxylated fatty acid glycerides, ester of fatty acids with polyols such as for example pentaerythrite or trimethylolpropane, fatty alcohol ethoxylates with narrow distribution of homologous or alkyl oligoglucosides as well as electrolytes such as sodium chloride and ammonium chloride.

Suitable **cationic polymers** are for example cationic cellulose derivatives, 15 such as e.g. a quaternized hydroxyethyl cellulose, which is available under the name of Polymer JR 400® from Amerchol, cationic starch, copolymers of diallyl ammonium salts and acrylamides, quaternized vinylpyrrolidone/vinylimidazol polymers, such as e.g. Luviquat® (BASF), condensation products of polyglycols and amines, quaternized collagen polypeptides, such as for example lauryl 20 dimonium hydroxypropyl hydrolyzed collagen (Lamequat® L / Grünau), quaternized wheat polypeptides, polyethyleneimine, cationic silicone polymers, such as e.g. amidomethicones, copolymers of adipic acid and dimethylamino hydroxypropyl diethylenetriamine (Cartaretine® / Sandoz), copolymers of acrylic acid with dimethyl diallylammonium chloride (Merquat® 550 /Chemviron), polyamino 25 polyamides, such as e.g. described in FR 2252840 A, as well as their cross-linked water soluble polymers, cationic chitin derivatives such as for example quaternized chitosan, possibly micro crystalline distributed, condensation products of dihalogen alkyls, such as e.g. dibromobutane with bisdialkylamines, such as e.g. bis-dimethylamino-1,3-propane, cationic guar-gum, such as e.g. Jaguar® 30 CBS, Jaguar® C-17, Jaguar® C-16 from Celanese, quaternised ammonium salt polymers, such as e.g. Mirapol® A-15, Mirapol® AD-1, Mirapol® AZ-1 from Miranol.

As exemplary **anionic, zwitterionic, amphoteric and non-ionic polymers** the following can be used: Vinyl acetate/crotonic acid copolymers, vinyl pyrrolidone/vinyl acrylate copolymers, vinyl acetate/butyl maleate/isobornyl

acrylate copolymers, methyl vinylether/maleic acid anhydride copolymers and their esters, non-cross-linked and with polyols cross-linked polyacrylic acids, acrylamido propyltrimethyl ammonium chloride/acrylate copolymers, octylacrylamide/methyl methacrylate/ tert.-butylaminoethyl methacrylate/2-5 hydroxypropyl methacrylate copolymers, polyvinylpyrrolidone, vinylpyrrolidone/vinylacetate copolymers, vinylpyrrolidone/ dimethylamino ethylmethacrylate/vinyl caprolactam terpolymers as well as possibly derivatized cellulose ethers and silicones.

10 Suitable **silicon compounds** are for example dimethyl polysiloxane, methylphenyl polysiloxane, cyclic silicones as well as amino, fatty acid, alcohol, polyether, epoxy, fluorine, glycoside and/or alkyl modified silicon compounds, which at room temperature can be in the liquid as well as in the resin state. Further suitable are simethicones, which are mixtures of dimethicones with an average chain length of 200 to 300 dimethyl siloxane units and hydrogenated silicates. A 15 detailed survey of suitable volatile silicones can also be found in Todd et al., *Cosm. Toil.* 91, 27 (1976).

20 Typical exemplary **fats** are glycerides, and as **waxes** natural waxes among others, can be used, such as e.g. candelilla wax, carnauba wax, Japan wax, espartogras wax, cork wax, guaruma wax, rice seed oil wax, sugar cane wax, ouricury wax, montan wax, beeswax, schellak wax, spermaceti, lanolin (wool wax), bürzel fat, ceresin, ozokerit (terrestrial wax), petrolatum, paraffin waxes, micro waxes; chemically modified waxes (hard waxes), such as e.g. montanester waxes, sasot waxes, hydrogenated yoyoba waxes as well as synthetic waxes, such as e.g. polyalkylene waxes and polyethylene glycol waxes.

25 As **stabilizers** metal salts of fatty acids, such as e.g. magnesium, aluminium and/or zinc stearate or ricinoleate can be used.

30 As **biogenic active substances** should be understood for example tocopherol, tocopherol acetate, tocopherol palmitate, ascorbic acid, desoxyribonucleic acid, retinol, bisabolol, allantoin, phytantriol, panthenol, AHA acids, aminoacids, ceramides, pseudoceramides, essential oils, extracts of plants and vitamin complexes.

As **deo active agents** e.g. antiperspirants such as aluminium chlorohydrate come into question. This agent is in the form of colourless, hygroscopic crystals, which easily melt in air, and is obtained through evaporation

of solutions of aluminium chloride in water. Aluminium chlorohydrate is used for manufacturing of perspiration inhibiting and deodorising preparations and has probably its effect through the partial closure of the perspiratory gland by means of precipitation of proteins and/or polysaccharides [see *J. Soc. Cosm. Chem.* 24, 5 281 (1973)]. Under the trade name Locron<sup>®</sup> of Hoechst AG, Frankfurt/FRG, an aluminium chlorohydrate is for example on the market, which corresponds to the formula  $[\text{Al}_2(\text{OH})_5\text{Cl}] \cdot 2.5 \text{ H}_2\text{O}$ , and use of this is especially preferred (see *J. Pharm. Pharmacol.* 26, 531 (1975)]. In addition to the chlorohydrates also aluminium hydroxylactates as well as acid aluminium/zirconium salts can be used.

10 As further deo active agents esterase inhibitors can be added. These are preferably trialkyl citrates such as trimethyl citrate, tripropyl citrate, triisopropyl citrate, tributyl citrate and especially triethyl citrate (Hydagen<sup>®</sup> CAT, Henkel KGaA, Düsseldorf/FRG). The substances inhibit the enzyme activity and thereby reduce the formation of odours. Probably the free acid is thereby set free through the

15 cleavage of the citric acid ester, and this acid lowers the pH value of the skin so much that the enzymes thereby are inhibited. Further substances which can be used as esterase inhibitors are sterol sulphates or phosphates, such as for example lanosterol, cholesterol, campesterol, stigmasterol and sitosterol sulphate or phosphate, Dicarboxylic acids and their esters, such as for example glutaric acid, glutaric acid monoethylester, glutaric acid diethylester, adipic acid, adipic acid monoethylester, adipic acid diethylester, malonic acid and malonic acid diethylester, hydroxycarboxylic acids and their esters, such as for example citric acid, malic acid, tartaric acid or tartaric acid diethylester. Antibacterial active

20 substances, which influence the germ flora and kill sweat destroying bacteria or inhibit their growth, can also be contained in the pin preparations. Examples of this are chitosan, phenoxyethanol and chlorohexidin gluconate. Also 5-chloro-2-(2,4-dichlorophen-oxy)-phenol has shown to have an especially good effect, and this product is marketed under the trade name Irgasan<sup>®</sup> by Ciba-Geigy, Basel/CH.

25

As **anti dandruff** agents climbazol, octopirox and zinc pyrethion can be used. Useable **film formation** agents are for example chitosan, microcrystalline chitosan, quaternary chitosan, polyvinylpyrrolidon, vinylpyrrolidon/vinylacetate copolymers, polymers of the acrylic acids, quaternary derivatives of cellulose, collagen, hyaluronic acid or its salts and similar compounds. As **swelling agents** for aqueous phases montmorillonite, clay mineral substances, pemulen, as well

as alkylmodified Carbopol types (Goodrich) can be used. Further suitable polymers or swelling agents can be found in the survey of R.Lochhead in *Cosm. Toil.* 108, 95 (1993).

**UV light protection factors** are e.g organic substances (light protection

5 filters) which by room temperature are in liquid or crystalline form, and which are capable of absorbing ultraviolet radiation and to set free the received energy in the form of radiation with long wavelength, e.g. in the form of heat. UVB filters can be soluble in oils or in water. As oil soluble substances the following are mentioned as examples:

10 • 3-Benzyliden camphor, respectively 3-benzylidene norcamphor and the derivatives thereof, e.g. 3-(4-methylbenzylidene) camphor as described in EP-B1 0693471;

15 • 4-aminobenzoic acid derivatives, preferably 4-(dimethylamino) benzoic acid 2-ethylhexylester, 4-(dimethylamino) benzoic acid 2-octylester and 4-(dimethylamino) benzoic acid amylester;

20 • esters of cinnamonic acid, preferably 4-methoxy cinnamonic acid 2-ethylhexylester, 4-methoxy cinnamonic acid propylester, 4-methoxy cinnamonic acid isoamylester, 2-cyano-3,3-phenyl cinnamonic acid 2-ethyhexylester (octocrylene);

25 • esters of salicylic acid, preferably salicylic acid 2-ethylhexylester, salicylic acid 4-isopropyl benzylester, salicylic acid homomenthylester;

• derivatives of benzophenone, preferably 2-hydroxy-4-methoxy benzophenone, 2-hydroxy-4-methoxy-4'-methyl benzophenone, 2,2'-dihydroxy-4-methoxy benzophenone;

30 • esters of benzalmalonic acid, preferably 4-methoxy benzmalonic acid 2-ethylhexyl ester,

• triazine derivatives, such as e.g. 2,4,6-trianilino-(p-carbo-2'-ethyl-1'-hexyloxy)-1,3,5-triazine and octyltriazone, as described in EP A1 0818450;

• propane-1,3-diones, such as e.g. 1-(4-tert.-butylphenyl)-3-(4'-methoxy-phenyl)-propane-1,3-dion;

35 • ketotricyclo(5,2,1,0)-decane derivatives, as described in EP-B1 06945521.

As water soluble substances the following can be mentioned:

• 2-Phenylbenzimidazol-5-sulphonic acid and the alkali, alkaline earth, ammonium, alkylammonium, alkanolammonium and glucammonium salts;

- sulphonic acid derivatives of benzophenones, preferably 2-hydroxy-4-methoxybenzophenon-5-sulphonic acid and their salts;
- sulphonic acid derivatives of 3-benzylidencamphen, such as e.g. 4-(2-oxo-3-bornylidenmethyl)-benzene sulphonic acid and 2-methyl-5-(2-oxo-bornyliden) sulphonic acid and their salts.

5 As typical UV-A filters especially derivatives of benzoyl methane comes in question, such as e.g. 1-(4'-tert.-butylphenyl)-3-(4'-methoxyphenyl)propane-1,3-dion, 4-tert.butyl-4'-methoxydibenzoyl-methane (Parsol 1789), or 1-phenyl-3-(4'-isopropylphenyl)-propane-1,3-dion. The UV-A and UV-B filters can of course also 10 be used in mixtures. In this case combinations of octocrylene or camphor derivatives with butyl methoxydibenzoylmethane are especially photosensitive.

15 In addition to the mentioned soluble substances also insoluble light protection pigments can be used for this purpose, i.e. fine disperse metal oxides or salts. Examples of suitable metal oxides are especially zinc oxide and titanium dioxide and in addition other oxides of iron, zirconium, silicon, manganese, aluminium and cerium, as well as their mixtures. As salts silicates (talk), barium sulphate or zinc stearate can be used. The oxides and salts are used in the form 20 of the pigments for skin caring and skin protecting emulsions and decorative cosmetics. The particles should have an average diameter of less than 100 nm, preferably between 5 and 50 nm and especially between 15 and 30 nm. They can have a spherical shape, but particles can also be used which have an ellipsoidal form or else have a shape which differs from the spherical shape. In sun 25 protecting agents preferably so-called micro or nano pigments are used. Preferably micronized zinc oxide is used. Further suitable UV light protection factors can be found in the survey by P.Finkel in *SÖFW-Journal 122, 543 (1996)*.

30 In addition to the primary light protection substances also secondary light protection substances of the **antioxidant** type find use, which interrupt the photochemical reaction chain, which is initiated when UV radiation penetrates the skin. Typical examples of such are amino acids (e.g. glycine, histidine, tyrosine, tryptophan) and their derivatives, imidazoles (e.g. urocanic acid) and their derivatives, peptides such as D,L-carnosine, D-carnosine, L-carnosine and their derivatives (e.g. anserine), carotenoids, carotene (e.g.  $\alpha$ -carotene,  $\beta$ -carotene, lycopene) and their derivatives, chlorogenic acid and its derivatives, liponic acid and its derivatives (e.g. dihydroliponic acid), aurothioglucose, propylthiouracil and

other thiols (e.g. thioredoxin, glutathion, cystein, cystin, cystamine and their glycosyl, n-acetyl, methyl, ethyl, propyl, amyl, butyl and lauryl, palmitoyl, oleyl,  $\gamma$ -linoleyl, cholesteryl and glyceryl esters) as well as their salts, dilauryl thiodipropionate, distearyl thiodipropionate, thiodipropionic acid and their derivatives (esters, ethers, peptides, lipides, nucleotides, nucleosides and salts) as well as sulfoximine compounds (e.g. buthionin sulfoximines, homocystein sulfoximines, butionin sulfones, penta-, hexa-, hepta-thionin sulfoximine) in very small compatible doses (e.g. pmol to  $\mu$ mol/kg), further (metal) chelating agents (e.g.  $\alpha$ -hydroxy fatty acids, palmitic acid, phytinic acid, lactoferrine),  $\alpha$ -hydroxy acids (e.g. citric acid, lactic acid, malic acid), humin acid, gallic acid, gallic extracts, bilirubin, bifiverdin, EDTA, EGTA and their derivatives, unsaturated fatty acids and their derivatives (e.g.  $\gamma$ -linolenic acid, linolic acid, oleic acid), folic acid and their derivatives, ubichinon and ubichinol and their derivatives, vitamin C and derivatives (e.g. ascorbyl palmitate, Mg-ascorbyl phosphate, ascorbyl acetate), tocopheroles and derivatives (e.g. vitamin E acetate), vitamin A and derivatives (vitamin A palmitate) as well as koniferyl benzoate of benzoe resin, rutinic acid and their derivatives,  $\alpha$ -glycosylrutin, ferula acid, furfuryliden glucitol, carnosine, butylhydroxy toluene, butylhydroxy anisol, nordihydro guajak resin acid, nordihydro guajaret acid, trihydroxy butyrophenon, uric acid and their derivatives, 20 mannose and its derivatives, super oxide dismutase, zinc and its derivatives (e.g. ZnO, ZnSO<sub>4</sub>), selen and its derivatives (e.g. selen-methionin), stilbenes and their derivatives (e.g. stilben oxide, trans-stilben oxide) and the derivatives suitable according to the invention (salts, esters, ethers, sugars, nucleotides, nucleosides, peptides and lipids) of these mentioned active substances.

25 For improvement of the flow properties further **hydrotropes**, such as for example ethanol, isopropyl alcohol, or polyols can be used. Polyols which in this case can be used preferably have 2 to 15 carbon atoms and at least two hydroxyl groups. The polyols can further contain additional functional groups, especially amino groups, or be modified with nitrogen. Typical examples are:

30

- Glycerol;
- alkylene glycols, such as for example ethylene glycol, diethylene glycol, propylene glycol, butylene glycol, hexylene glycol as well as polyethylene glycols with an average molecular weight from 100 to 1 000 Daltons;

- oligoglycerol mixtures of technical quality with a self-condensation degree of 1.5 to 10, such as e.g. technical quality diglycerol mixtures with a diglycerol content of 40 to 50 % by weight;
- methyol compounds, such as especially trimethylol ethane, trimethylol propane, trimethylol butane, pentaerythrite and dipentaerythrite;
- low alkyl glucosides, especially such with 1 to 8 carbons in the alkyl residue, such as for example methyl and butyl glucoside;
- sugar alcohols with 5 to 12 carbon atoms, such as for example sorbitol or mannit;
- sugars with 5 to 12 carbon atoms, such as for example glucose or saccharose;
- aminosugars, such as for example glucamine;
- dialcoholamines, such as diethanolamine or 2-amino-1,3-propanediol.

As **preservatives** for example phenoxyethanol, formaldehyde solution, parabene, pentanediol or sorbic acid as well as those mentioned in enclosure 6, parts A and B of the cosmetic regulation, are further classes of substances. As **insect repellents** N,N-diethyl-m-toluamide, 1,2-pantanediol or insect repellent 3535 come into question, as **self tanning agent** dihydroxyacetone is suited.

As **perfume oils** mixtures of natural and synthetic scent substances should be mentioned. Natural scent substances are extracts of flowers (lilies, lavendel, roses, jasmin, neroli, ylang-ylang), stems and blades (geranium, patchouli, petitgrain), fruits (anis, coriander, caraway, juniper), fruit shells (bergamot, lemon, orange), roots (macis, angelica, celery, kardamon, costus, iris, calmus), wood (stone pine, sandel, guajac, cedar, rosewood), herbs and grass (tarragon, lemongrass, sage, thyme), needles and twigs (spruce, fir, pine, traipsed), resins and balsams (galbanum, elemi, benzoe, myrrh, olibanum, opopanax). Raw materials from animals are also possible, such as for example zibet and castoreum. Typical synthetic odour compounds are products from types of esters, ethers, aldehydes, ketones, alcohols and hydrocarbons. Odour compounds from types of esters are e.g. benzyl acetate, phenoxyethyl isobutyrate, p-tert.-butylcyclohexyl acetate, linalyl acetate, dimethylbenzylcarbinyl acetate, phenylethyl acetate, linalyl benzoate, benzyl formate, ethylmethylphenyl glycinate, allylcyclohexyl propionate, styrallyl propionate and benzyl salicylate. Benzylethyl ether belongs for example to the ethers, to the aldehydes e.g. the linear alkanals

with 8 to 18 carbon atoms, citral, citronellal, citronellyl oxyacetaldehyde, cyclamen aldehyde, hydroxy citronellal, linalol and bourgeonal, to the ketones e.g. the ionones,  $\alpha$ -isomethyl ionon and methylcedryl ketone, to the alcohols anethol, citronellol, eugenol, isoeugenol, geraniol, linalool, phenylethyl alcohol and 5 terpineol; to the hydrocarbons mainly the terpenes and balsams belong. However, mixtures of different odour substances are preferred, which together give a pleasant smell. Also etheral oils with low volatility, which often are used as aroma components, are suited as perfume oils, e.g. sage oil, chamomile oil, carnation oil, melissa oil, mint oil, cinnamon leaf oil, limeflower oil, juniper berry oil, vetiver oil, 10 oliban oil, galbanum oil, labolanum oil and lavandin oil. Preferably used are bergamot oil, dihydromyrcenol, linalol, lyral, citronellol, phenylethyl alcohol,  $\alpha$ -hexylcinnamon aldehyde, geraniol, benzylacetone, cyclamen aldehyde, linalool, boisambrene forte, ambroxane, indol, hedione, sandelice, lemon oil, mandarin oil, orangenoil, allylamyl glycolate, cyclovertal, lavandine oil, muskateller sage oil, 15  $\beta$ -damascone, geranium oil bourbon, cyclohexyl salicylate, vertofix coeur, iso-E-super, fixolide NP, evermyl, iraidein gamma, phenylacetic acid, geranyl acetate, benzyl acetate, rose oxide, romillate, irotyl and floramate, alone or in mixtures.

As **colouring agents** such substances which are suited and approved for cosmetic purposes can be used, such as for example those mentioned in the 20 publication "Kosmetische Färbemittel" (cosmetic dyes) of the "Farbstoffkommission der Deutschen Forschungsgemeinschaft", published by Verlag Chemie, Weinheim, 1984, p. 81-106. These dyes are generally used in concentrations from 0.001 to 0.1 % by weight, based on the whole mixture.

Typical examples of **germ inhibiting** substances are preservatives with 25 specific effects against gram-positive bacteria, such as 2,4,4'-trichloro-2'-hydroxy diphenylether, chlorohexidin (1,6-di-(4-chlorophenyl-biguanido-hexan) or TCC (3,4,4'-trichlorocarbanilide). Many scent substances and etheral oils also have antimicrobial properties. Typical examples are the active agents eugenol, menthol and thymol in carnation, mint and thyme oil. An interesting natural deo substance 30 is the terpene alcohol famesol (3,7,11-trimethyl-2,6,10-dodecatrien-1-ol), which is present in lime flower oil and has a smell of lilies of the valley. Also glycerol monolaurate have been used as bacteriostaticum. Normally the content of the further germ inhibiting agent is about 0.1 to 2 % by weight - based on the solids content of the preparations.

The cumulative contents of the auxiliary and additional agents can be 1 to 50, preferably 5 to 40 % by weight, based on the agents. The manufacture of the agents can take place by common cold or hot processes; preferably the work is carried out according to the phase inversion temperature method.

5

## Examples

A panel consisting of 15 female probands aged between 35 and 50 years were during a time period of 28 days daily exposed to a daily exposition of different glucans and/or chitosans. The probands used the skin cremes daily before going to bed. With intervals of 7 days the number, depth and lenght of the skin wrinkles were determined for each of the participants by means of profilometry of a selected part of the skin, i.e. a vertical stripe of 2 cm width and 5 cm length, having an upper left and right boundary. which occurs if from the nose root a horizontal line is drawn, from this and against the right eye 2, respectively 4 cm, are cleared away and both resulting points in each case are elongated in an angle of 270° in each case 2 cm. The dimensionless product of depth, number and lenght of the skin wrinkles on the day before the beginning of the exposure was set as standard (= 100 %), and all the following measurements were based on this. At the same time the skin roughness of the pro-bands was evaluated on a scale from 0 = "unchanged" to 3 = "strongly improved". The results are summarized in Table 1. Examples 1 and 2 are according to the invention, the examples V1 to V3 are for comparison.

25

**Table 1**  
**Skin ageing and skin roughness**

<b>Composition / Performance</b>	<b>1</b>	<b>2</b>	<b>V1</b>	<b>V2</b>	<b>V3</b>
Cetyl stearyl alcohol	8.0	8.0	8.0	8.0	8.0
Ceteareth-12	1.5	1.5	1.5	1.5	1.5
Ceteareth-20	1.5	1.5	1.5	1.5	1.5
Cetearyl isononanoate	15.0	15.0	15.0	15.0	15.0
Paraffin oil, viscous	5.0	5.0	5.0	5.0	5.0
Baysilon oil M 300	5.0	5.0	5.0	5.0	5.0
$\beta$ -1,3 Glucan *	20.0	20.0	20.0	-	-
Chitosan **	2.0	-	-	2.0	-
Succinilated chitosan ***	-	2.0	-	-	2.0
Glycerol	6.0	6.0	6.0	6.0	6.0
Water	ad 100				
<b><i>Skin ageing [%-rel]</i></b>					
- bevor the treatment	100	100	100	100	100
- after 7 d	91	92	96	99	99
- after 14 d	85	87	91	97	97
- after 21 d	80	83	85	95	95
- after 28 d	73	75	79	91	91
<b><i>Skin roughness</i></b>					
- bevor the treatment	0	0	0	0	0
- after 7 d	2	1	1	0	0
- after 14 d	3	2	2	0	0
- after 21 d	3	3	3	1	1
- after 28 d	3	3	3	1	1

\*) Highcareene® GS

\*\*) Hydagen® CMF

\*\*\*) Hydagen® SCD (all are from Henkel KGaA, Düsseldorf/FRG)

The following table contains formulation examples.

**Table 1 - Cosmetic Preparations (water, preservatives ad 100 % by weigh)**

Composition (INCI)	1	2	3	4	5	6	7	8	9	10
<b>Texapon® NSO</b> Sodium latureth. sulphate	-	-	-	-	-	-	38.0	38.0	25.0	-
<b>Texapon® SB 3</b> Disodium laureth. sulphosuccinate	-	-	-	-	-	-	-	-	10.0	-
<b>Plantacare® 818</b> Coco glucosides	-	-	-	-	-	-	7.0	7.0	6.0	-
<b>Plantacare® PS 10</b> Sodium laureth.sulphate (and) coco glucosides	-	-	-	-	-	-	-	-	-	16.0
<b>Dehyton® PK 45</b> Cocamidopropyl betaine	-	-	-	-	-	-	-	-	10.0	-
<b>Dehyquart® A</b> Centrimonium chloride	2.0	2.0	2.0	2.0	4.0	4.0	-	-	-	-
<b>Dehyquart L® 80</b> Dicocoylmethylethoxymonium methosulphate (and) propylene glycol	1.2	1.2	1.2	1.2	0.6	0.6	-	-	-	-
<b>Eumulgin® B2</b> Ceteareth-20	0.8	0.8	-	0.8	-	1.0	-	-	-	-
<b>Eumulgin® VL 75</b> Lauryl glucoside (and) polyglyceryl-2 polyhydroxy stearate (and) glycerol	-	-	0.8	-	0.8	-	-	-	-	-
<b>Lanette® O</b> Cetearyl alcohol	2.5	2.5	2.5	2.5	3.0	2.5	-	-	-	-
<b>Cutina® GMS</b> Glyceryl stearate	0.5	0.5	0.5	0.5	0.5	1.0	-	-	-	-
<b>Cetiol® HE</b> PEG-7 glyceryl cocoate	1.0	-	-	-	-	-	-	-	1.0	-
<b>Cetiol® PGL</b> Hexyldecanol (and) hexyldecyll laurate	-	1.0	-	-	1.0	-	-	-	-	-
<b>Cetiol® V</b> Decyl oleate	-	-	-	1.0	-	-	-	-	-	-
<b>Eutanol® G</b> Octyldodecanol	-	-	1.0	-	-	1.0	-	-	-	-
<b>Nutrilan® Keratin W</b> Hydrolyzed keratine	-	-	-	2.0	-	-	-	-	-	-
<b>Lamesoft® LMG</b> Glyceryl laurate (and) potassium cocoyl hydrolyzed collagen	-	-	-	-	-	-	3.0	2.0	4.0	-
<b>Euperlan® PK 3000 AM</b> Glyceryl distearate (and) laureth.-4 (and) cocamidopropyl betaine	-	-	-	-	-	-	-	3.0	5.0	5.0
<b>Generol® 122 N</b> Soya sterol	-	-	-	-	1.0	1.0	-	-	-	-
<b>Highcareen® GS</b> Betaglucan	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
<b>Desoxy ribonucleic acid</b> Molecular weight approx. 70000	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2
<b>Copherol® 12250</b> Tocopherol acetate	-	-	0.1	0.1	-	-	-	-	-	-
<b>Artypon® F</b> Laureth-2	-	-	-	-	-	-	3.0	3.0	1.0	-
<b>Sodium chloride</b>	-	-	-	-	-	-	-	1.5	-	1.5

(1-4) Hair rinsing, (5-6) Hair cure, (7-8) Shower bath, (9) Shower gel, (10) Cleaning lotion

**Table 1 Cosmetic preparations (water, preservatives ad 100 % by weight) - (cont.)**

Composition (INCI)	11	12	13	14	15	16	17	18	19	20
<b>Texapon® NSO</b> Sodium laureth. sulphate	20.0	20.0	12.4	-	25.0	11.0	-	-	-	-
<b>Texapon® K 14 S</b> Sodium myreth. sulphate	-	-	-	-	-	-	-	-	11.0	23.0
<b>Texapon® SB 3</b> Disodium laureth. sulphosuccinate	-	-	-	-	-	7.0	-	-	-	-
<b>Plantacare® 818</b> Coco glucosides	5.0	5.0	4.0	-	-	-	-	-	6.0	4.0
<b>Plantacare® 2000</b> Decyl glucoside	-	-	-	-	5.0	4.0	-	-	-	-
<b>Plantacare® PS 10</b> Sodium laureth. sulphate (and) coco glucosides	-	-	-	40.0	-	-	16.0	17.0	-	-
<b>Dehyton® PK 45</b> Cocamidopropyl betaine	20.0	20.0	-	-	8.0	-	-	-	-	7.0
<b>Eumulgin® B2</b> Ceteareth-20	-	-	-	1.0	1.0	-	-	-	-	-
<b>Lameform® TGI</b> Polyglyceryl-3 isostearate	-	-	-	4.0	-	-	-	-	-	-
<b>Dehymuls® PGPH</b> Polyglyceryl-2 dipolyhydroxy stearate	-	-	1.0	-	-	-	-	-	-	-
<b>Monomuls® 90-L 12</b> Glyceryl laurate	-	-	-	-	-	-	-	-	-	1.0
<b>Cutina® GMS</b> Glyceryl stearate	-	-	-	-	-	-	-	-	1.0	-
<b>Cetiol® HE</b> PEG-7 Glyceryl cocoate	-	0.2	-	-	-	-	-	-	-	-
<b>Eutanol® G</b> Octyldodecanol	-	-	-	3.0	-	-	-	-	-	-
<b>Nutrilan® Keratin W</b> Hydrolyzed keratin	-	-	-	-	-	-	-	-	2.0	2.0
<b>Nutrilan® I</b> Hydrolyzed collagen	1.0	-	-	-	-	2.0	-	2.0	-	-
<b>Lamesoft® LMG</b> Glyceryl laurate (and) potassium cocoyl hydrolyzed collagen	-	-	-	-	-	-	-	-	1.0	-
<b>Lamesoft® 156</b> Hydrogenated tallow glyceride (and) potassium cocoyl hydrolyzed collagen	-	-	-	-	-	-	-	-	-	5.0
<b>Gluadin® WK</b> Sodium cocoyl hydrolyzed wheat protein	1.0	1.5	4.0	1.0	3.0	1.0	2.0	2.0	2.0	-
<b>Euperlan® PK 3000 AM</b> Glycol distearate (and) laureth-4 (and) cocamidopropyl betaine	5.0	3.0	4.0	-	-	-	-	3.0	3.0	-
<b>Panthenol</b>	-	-	1.0	-	-	-	-	-	-	-
<b>Highcareen® GS</b> Betaglucan	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
<b>Desoxy ribonucleic acid</b> Molecular weight approx. 70000	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2
<b>Arlypon® F</b> Laureth-2	2.6	1.6	-	1.0	1.5	-	-	-	-	-
<b>Sodium chloride</b>	-	-	-	-	-	1.6	2.0	2.2	-	3.0
<b>Glycerol (86 % by weight)</b>	-	5.0	-	-	-	-	-	1.0	3.0	-

(11-14) Shower bath "two-in-one". (15-20) Shampoo

**Table 1 - Cosmetic preparations (water, preservatives ad 100 % by weight) - (cont. 2)**

Composition (INCI)	21	22.	23	24	25	26	27	28	29	30
<b>Texapon® NSO</b> Sodium laureth. sulphate	-	30.0	30.0	-	25.0	-	-	-	-	-
<b>Plantacare® 818</b> Coco glucosides	-	10.0	-	-	20.0	-	-	-	-	-
<b>Plantacare® PS 10</b> Sodium laureth. sulphate (and) coco glucosides	22.0	-	5.0	22.0	-	-	-	-	-	-
<b>Dehyton® PK 45</b> Cocamidopropyl betaine	15.0	10.0	15.0	15.0	20.0	-	-	-	-	-
<b>Emulgade® SE</b> Glyceryl stearate (and) ceteareth. 12/20 (and) cetearyl alcohol (and) cetyl palmitate	-	-	-	-	-	5.0	5.0	4.0	-	-
<b>Eumulgin® B1</b> Ceteareth-12	-	-	-	-	-	-	-	1.0	-	-
<b>Lameform® TGI</b> Polyglyceryl-3 isostearate	-	-	-	-	-	-	-	-	4.0	-
<b>Dehymuls® PGPH</b> Polyglyceryl-2 dipolyhydroxystearate	-	-	-	-	-	-	-	-	-	4.0
<b>Monomuls® 90-O 18</b> Glyceryl oleate	-	-	-	-	-	-	-	-	2.0	-
<b>Cetiol® HE</b> PEG-7 Glyceryl cocoate	2.0	-	-	2.0	5.0	-	-	-	-	2.0
<b>Cetiol® OE</b> Dicaprylyl ether	-	-	-	-	-	-	-	-	5.0	6.0
<b>Cetiol® PGL</b> Hexyldecanol (and) hexyldecyll laurate	-	-	-	-	-	-	-	3.0	10.0	9.0
<b>Cetiol® SN</b> Cetearyl isononanoate	-	-	-	-	-	3.0	3.0	-	-	-
<b>Cetiol® V</b> Decyl oleate	-	-	-	-	-	3.0	3.0	-	-	-
<b>Myritol® 318</b> Coco caprylate caprate	-	-	-	-	-	-	-	3.0	5.0	5.0
<b>Bees Wax</b>	-	-	-	-	-	-	-	-	7.0	5.0
<b>Nutrilan® Elastin E20</b> Hydrolyzed elastin	-	-	-	-	-	2.0	-	-	-	-
<b>Nutrilan® I-50</b> Hydrolyzed collagen	-	-	-	-	2.0	-	2.0	-	-	-
<b>Gluadin® AGP</b> Hydrolyzed wheat glutene	0.5	0.5	0.5	-	-	-	-	0.5	-	-
<b>Gluadin® WK</b> Sodium cocoyl hydrolyzed wheat protein	2.0	2.0	2.0	2.0	5.0	-	-	-	0.5	0.5
<b>Eupertan® PK 3000 AM</b> Glycol distearate (and) laureth-4 (and) cocamidopropyl betaine	5.0	-	-	5.0	-	-	-	-	-	-
<b>Highcareen® GS</b> Betaglucan	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
<b>Desoxy ribonucleic acid</b> Molecular weight approx. 70000	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2
<b>Magnesium sulphate heptahydrate</b>	-	-	-	-	-	-	-	-	1.0	1.0
<b>Glycerol (86 % by weight)</b>	-	-	-	-	-	3.0	3.0	5.0	5.0	3.0

(21-25) Foam bath, (26) Soft creme, (27.28) Moisture emulsion, (29.30) Night creme

**Table 1 - Cosmetic preparations (water, preservatives ad 100 % by weight) - (cont. 3)**

Composition (INCI)	31	32	33	34	35	36	37	38	39	40
<b>Dehymuls® PGPH</b>	4.0	3.0	-	5.0	-	-	-	-	-	-
Polyglyceryl-2 dipolyhydroxystearate										
<b>Lameform® TGI</b>	2.0	1.0	-	-	-	-	-	-	-	-
Polyglyceryl-3 diisostearate										
<b>Emulgade® PL 68/50</b>	-	-	-	-	4.0	-	-	-	3.0	-
Cetearyl glucoside (and) cetearyl alcohol										
<b>Eumulgin® B2</b>	-	-	-	-	-	-	-	2.0	-	-
Ceteareth-20										
<b>Tegocare® PS</b>	-	-	3.0	-	-	-	4.0	-	-	-
Polyglyceryl-3 methylglucose distearate										
<b>Eumulgin VL75</b>	-	-	-	-	-	3.5	-	-	2.5	-
Polyglyceryl-2 dipolyhydroxystearate (and) lauryl glucoside (and) glycerol										
<b>Beeswax</b>	3.0	2.0	5.0	2.0				-		
<b>Cutina® GMS</b>	-	-	-	-	-	2.0	4.0	-	-	4.0
Glyceryl stearate										
<b>Lanette® O</b>	-	-	2.0	-	2.0	4.0	2.0	4.0	4.0	1.0
Cetearyl alcohol										
<b>Antaron® V 216</b>	-	-	-	-	-	3.0	-	-	-	2.0
PVP / hexadecene copolymer										
<b>Myritol® 818</b>	5.0	-	10.0	-	8.0	6.0	6.0	-	5.0	5.0
Coco glycerides										
<b>Finsolv® TN</b>	-	6.0	-	2.0	-	-	3.0		-	2.0
C12/15 Alkyl benzoate										
<b>Cetiol® J 600</b>	7.0	4.0	3.0	5.0	4.0	3.0	3.0	-	5.0	4.0
Oleyl erucate										
<b>Cetiol® OE</b>	3.0	-	6.0	8.0	6.0	5.0	4.0	3.0	4.0	6.0
Dicaprylyl ether										
<b>Mineral Oil</b>	-	4.0	-	4.0	-	2.0	-	1.0	-	-
<b>Cetiol® PGL</b>	-	7.0	3.0	7.0	4.0	-	-	-	1.0	-
Hexadecanol (and) hexyl laurate										
<b>Panthenol / Bisabolol</b>	1.2	1.2	1.2	1.2	1.2	1.2	1.2	1.2	1.2	1.2
<b>Highcareen® GS</b>	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Betaglucan										
<b>Desoxy ribonucleic acid</b>	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2
Molecular weight approx. 70000										
<b>Copherol® F 1300</b>	0.5	1.0	1.0	2.0	1.0	1.0	1.0	2.0	0.5	2.0
Tocopherol / tocopheryl acetate										
<b>Neo Heliopan® Hydro</b>	3.0	-	-	3.0	-	-	2.0	-	2.0	-
Sodium phenylbenzimidazole sulphonate										
<b>Neo Heliopan® 303</b>	-	5.0	-	-	-	4.0	5.0	-	-	10.0
Octocrylene										
<b>Neo Heliopan® BB</b>	1.5	-	-	2.0	1.5	-	-	-	2.0	-
Benzophenone-3										
<b>Neo Heliopan® E 1000</b>	5.0	-	4.0	-	2.0	2.0	4.0	10.0	-	-
Isoamyl p-methoxycinnamate										
<b>Neo Heliopan® AV</b>	4.0	-	4.0	3.0	2.0	3.0	4.0	-	10.0	2.0
Octyl methoxycinnamate										
<b>Uvinul® T 150</b>	2.0	4.0	3.0	1.0	1.0	1.0	4.0	3.0	3.0	3.0
Octyl triazone										
<b>Zinc oxide</b>	-	6.0	6.0	-	4.0	-	-	-	-	5.0
<b>Titanium dioxide</b>	-	-	-	-	-	-	-	5.0	-	-
<b>Glycerol (86 % by weight)</b>	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0

(31) W/O Sun protection creme, (32-34) W/O Sun protection lotion, (35, 38,40) O/W Sun protection lotion (36, 37, 39) O/W Sun protection creme

## P a t e n t c l a i m s

1. Cosmetic preparations, containing
  - (a) water soluble  $\beta$ -(1,3) glucans, substantially free from  $\beta$ -(1,6) linkages, and
  - 5 (b) chitosans.
2. Preparations according to claim 1, **characterised by** that they contain glucans which are obtained based on yeasts from the family *Saccharomyces*.
- 10 3. Preparations according to claim 1 and/or 2, **characterised by** that they contain glucans which are obtained by contacting glucans with  $\beta$ -(1,3) and  $\beta$ -(1,6) linkages with  $\beta$ -(1,6) glucanases, in such a way that practically all  $\beta$ -(1,6) linkages are loosened.
- 15 4. Preparations according to claim 3, **characterised by** that glucans are used, which previously have been treated with glucanases based on *Trichoderma harzianum*.
5. Preparations according to at least one of the claims 1 to 4, **characterised** 20 **by** that they contain chitosans with molecular weights in the area from 50 000 to 500 000 Daltons.
- 25 6. Preparations according to at least one of the claims 1 to 4, **characterised** **by** that they contain chitosans with molecular weights in the area from 800 000 to 1 200 000 Daltons.
7. Preparations according to at least one of the claims 1 to 6, **characterised** **by** that they contain carboxylated chitosans.
- 30 8. Preparations according to at least one of the claims 1 to 7, **characterised** **by** that they contain succinylated chitosans.
9. Preparations according to at least one of the claims 1 to 8, **characterised** **by** that they contain

(a) 0.01 to 25 % by weight of water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages, and

(b) 0.01 to 5 % by weight of chitosans,  
provided that the stated amounts are supplemented with water as well as  
5 optionally other auxiliaries and additional agents up 100 % by weight.

10. Use of mixtures containing

(a) water soluble  $\beta$ -(1,3) glucans, which are substantially free from  $\beta$ -(1,6) linkages,  
10 (b) chitosans,  
for manufacturing of cosmetic preparations.

## ABSTRACT

The invention relates to cosmetic preparations containing (a) water-soluble  $\beta$ -(1,3) glucans, substantially devoid of  $\beta$ -(1,6) links, and (b) chitosans. The agents are suitable for hair care and personal hygiene and can also be used for sun protection.

Docket: CU-2652

L 698184744

**COMBINED DECLARATION AND POWER OF ATTORNEY***(ORIGINAL, DESIGN, NATIONAL STAGE OF PCT, SUPPLEMENTAL, DIVISIONAL,  
CONTINUATION OR CIP)*

As a below named inventor, I hereby declare that:

**TYPE OF DECLARATION**

This declaration is of the following type: *(check one applicable item below)*

- original
- design
- supplemental

*Note: If the Declaration is for an International Application being filed as a divisional, continuation or continuation-in-part application, do not check next item; check appropriate one of last three items.*

- national stage of PCT

*Note: If one of the following 3 items apply, then complete and also attach ADDED PAGES FOR DIVISIONAL, CONTINUATION OR CIP.*

- divisional
- continuation
- continuation-in-part (CIP)

**INVENTORSHIP IDENTIFICATION**

*WARNING: If the inventors are each not the inventors of all the claims, an explanation of the facts, including the ownership of all the claims at the time the last claimed invention was made, should be submitted.*

My residence, post office address and citizenship are as stated below, next to my name. I believe that I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter that is claimed, and for which a patent is sought on the invention entitled:

**TITLE OF INVENTION**

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COSMETIC PREPARATIONS

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## **SPECIFICATION IDENTIFICATION**

the specification of which: (*complete (a), (b) or (c)*)

(a) is attached hereto.

(b) was filed on \_\_\_\_\_ as  Serial No. \_\_\_\_\_ or  
 Express Mail No. (as Serial No. not yet known) \_\_\_\_\_  
and was amended on \_\_\_\_\_ (if applicable).

**Note:** *Amendments filed after the original papers are deposited with the PTO that contain new matter are not accorded a filing date by being referred to in the Declaration. Accordingly, the amendments involved are those filed with the application papers or, in the case of a supplemental Declaration, are those amendments claiming matter not encompassed in the original statement of invention or claims. See 37 CFR 1.67.*

(c) was described and claimed in PCT International Application No. PCT/EP00/01837 filed on 03 March 2000.

**ACKNOWLEDGEMENT OF REVIEW OF PAPERS AND DUTY OF CANDOR**

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information, which is material to patentability as defined in 37, Code of Federal Regulations, § 1.56,

*(also check the following items, if desired)*

- and which is material to the examination of this application, namely, information where there is a substantial likelihood that a reasonable Examiner would consider it important in deciding whether to allow the application to issue as a patent, and
- in compliance with this duty, there is attached an information disclosure statement, in accordance with 37 CFR 1.98.

**PRIORITY CLAIM (35 U.S.C. § 119(a)-(d))**

I hereby claim foreign priority benefits under Title 35, United States Code, § 119(a)-(d) of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed.

(complete (d) or (e))

(d) no such applications have been filed.  
 (e) such applications have been filed as follows.

*Note: Where item (c) is entered above and the international application which designated the U.S. itself claimed priority check item (e), enter the details below and make the priority claim.*

**PRIOR FOREIGN/PCT APPLICATION(S) FILED WITHIN 12 MONTHS  
(6 MONTHS FOR DESIGN) PRIOR TO THIS APPLICATION  
AND ANY PRIORITY CLAIMS UNDER 35 U.S.C. § 119(a)-(d)**

COUNTRY (OR INDICATE IF PCT)	APPLICATION NUMBER	DATE OF FILING (day/month/year)	PRIORITY CLAIMED UNDER 35 USC 119
Germany	199 11 056.5	12 March 1999	<input checked="" type="checkbox"/> YES <input type="checkbox"/> NO
			<input type="checkbox"/> YES <input type="checkbox"/> NO

**CLAIM FOR BENEFIT OF PRIOR U.S. PROVISIONAL APPLICATION(S)  
(35 U.S.C. § 119(e))**

I hereby claim the benefit under Title 35, United States Code, § 119(e) of any United States provisional application(s) listed below:

PROVISIONAL APPLICATION NUMBER	FILING DATE

**ALL FOREIGN APPLICATION(S), IF ANY, FILED MORE THAN 12 MONTHS  
(6 MONTHS FOR DESIGN) PRIOR TO THIS U.S. APPLICATION**

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*Note: If the application filed more than 12 months from the filing date of this application is a PCT filing forming the basis for this application entering the United States as (1) the national stage or (2) a continuation, divisional, or continuation-in-part, then also complete ADDED PAGES TO COMBINED DECLARATION AND POWER OF ATTORNEY FOR DIVISIONAL, CONTINUATION OR CIP APPLICATION for benefit of the prior U.S. or PCT application(s) under 35 U.S.C. § 120.*

## POWER OF ATTORNEY

I hereby appoint the following practitioner(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith (*list name and registration number*).

Thomas F. Peterson, 24790; Richard J. Streit, 25765; Donald P. Reynolds, 26220; W. Dennis Drehkoff, 27193; Vangelis Economou, 32341; Brian W. Hameder, 45613; Valerie Neymeyer-Tynkov, 46956; Paul B. West, 18947; Joseph H. Handelman, 26179; Peter D. Galloway 27885; John Richards, 31503; Iain C. Baillie, 24090; Richard P. Berg, 28145 10

Attached, as part of this declaration and power of attorney, is the authorization of the above-named practitioner(s) to accept and follow instructions from my representative(s).

---

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---

## DECLARATION

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

## SIGNATURE(S)

*Note: Carefully indicate the family (or last) name, as it should appear on the filing receipt and all other documents.*

### Full name of first joint inventor

100  
Ute  
(Given Name)

*Ute*  
Middle Initial or Name

GRIESBACH  
(Family (or Last) Name)

### Inventor's signature

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2-00

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3-00

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4-00

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5-00

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**PATENT**

Docket: CU-2652

L 698 184744

**COMBINED DECLARATION AND POWER OF ATTORNEY***(ORIGINAL, DESIGN, NATIONAL STAGE OF PCT, SUPPLEMENTAL, DIVISIONAL,  
CONTINUATION OR CIP)*

As a below named inventor, I hereby declare that:

**TYPE OF DECLARATION**This declaration is of the following type: *(check one applicable item below)*

original  
 design  
 supplemental

*Note: If the Declaration is for an International Application being filed as a divisional, continuation or continuation-in-part application, do not check next item; check appropriate one of last three items.*

national stage of PCT

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divisional  
 continuation  
 continuation-in-part (CIP)

**INVENTORSHIP IDENTIFICATION**

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**TITLE OF INVENTION**

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COSMETIC PREPARATIONS

---

**SPECIFICATION IDENTIFICATION**

the specification of which: (complete (a), (b) or (c))

(a) is attached hereto.

(b) was filed on \_\_\_\_\_ as  Serial No. \_\_\_\_\_ or  
 Express Mail No. (as Serial No. not yet known) \_\_\_\_\_  
and was amended on \_\_\_\_\_ (if applicable).

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*(also check the following items, if desired)*

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in compliance with this duty, there is attached an information disclosure statement, in accordance with 37 CFR 1.98.

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I hereby claim foreign priority benefits under Title 35, United States Code, § 119(a)-(d) of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed.

(complete (d) or (e))

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**PRIOR FOREIGN/PCT APPLICATION(S) FILED WITHIN 12 MONTHS  
(6 MONTHS FOR DESIGN) PRIOR TO THIS APPLICATION  
AND ANY PRIORITY CLAIMS UNDER 35 U.S.C. § 119(a)-(d)**

COUNTRY (OR INDICATE IF PCT)	APPLICATION NUMBER	DATE OF FILING (day/month/year)	PRIORITY CLAIMED UNDER 35 USC 119
Germany	199 11 056.5	12 March 1999	<input checked="" type="checkbox"/> YES <input type="checkbox"/> NO
			<input type="checkbox"/> YES <input type="checkbox"/> NO

**CLAIM FOR BENEFIT OF PRIOR U.S. PROVISIONAL APPLICATION(S)  
(35 U.S.C. § 119(e))**

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**POWER OF ATTORNEY**

I hereby appoint the following practitioner(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith (*list name and registration number*).

Thomas F. Peterson, 24790; Richard J. Streit, 25765; Donald P. Reynolds, 26220; W. Dennis Drehkoff, 27193; Vangelis Economou, 32341; Brian W. Hameder, 45613; Valerie Neymeyer-Tynkov, 46956; Paul B. West, 18947; Joseph H. Handelman, 26179; Peter D. Galloway 27885; John Richards, 31503; Iain C. Baillie, 24090; Richard P. Berg, 28145

Attached, as part of this declaration and power of attorney, is the authorization of the above-named practitioner(s) to accept and follow instructions from my representative(s).

**SEND CORRESPONDENCE TO:**

Richard J. Streit  
c/o Ladas & Parry  
224 South Michigan Avenue  
Suite 1200  
Chicago, Illinois 60604

**DIRECT TELEPHONE CALLS TO:**

(*Name and telephone number*)

(312) 427-1300

**DECLARATION**

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

**SIGNATURE(S)**

*Note: Carefully indicate the family (or last) name, as it should appear on the filing receipt and all other documents.*

**Full name of first joint inventor**

Ute	(Given Name)		GRIESBACH
		(Middle Initial or Name)	(Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** \_\_\_\_\_ **Germany** \_\_\_\_\_

**Residence** \_\_\_\_\_ **Dusseldorf, Germany** \_\_\_\_\_

**Post Office Address** \_\_\_\_\_ **Ludolfstr. 13, D-40597 Dusseldorf, Germany** \_\_\_\_\_

**Full name of second joint inventor**

Rolf \_\_\_\_\_ WACHTER \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** \_\_\_\_\_ **Germany** \_\_\_\_\_

**Residence** \_\_\_\_\_ Dusseldorf, Germany \_\_\_\_\_

**Post Office Address** \_\_\_\_\_ Clausthal-Zellerfelder-Str. 48, D-40595 Dusseldorf, Germany \_\_\_\_\_

**Full name of third joint inventor**

Achim \_\_\_\_\_ ANSMANN \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** \_\_\_\_\_ **Germany** \_\_\_\_\_

**Residence** \_\_\_\_\_ Erkrath, Germany \_\_\_\_\_

**Post Office Address** \_\_\_\_\_ Kirchberg 25, D-40699 Erkrath, Germany \_\_\_\_\_

**Full name of fourth joint inventor**

Bernd \_\_\_\_\_ FABRY \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** 26 October 2001 **Country of Citizenship** \_\_\_\_\_ **Germany** \_\_\_\_\_

**Residence** \_\_\_\_\_ Korschenbroich, Germany \_\_\_\_\_

**Post Office Address** \_\_\_\_\_ Danziger Str. 31, D-41352 Korschenbroich, Germany \_\_\_\_\_

**Full name of fifth joint inventor**

Wolf \_\_\_\_\_ EISFELD \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** \_\_\_\_\_ **Germany** \_\_\_\_\_

**Residence** \_\_\_\_\_ Dusseldorf, Germany \_\_\_\_\_

**Post Office Address** \_\_\_\_\_ Am Monchgraben 4, D-40597 Dusseldorf, Germany \_\_\_\_\_

**Full name of sixth joint inventor**

Rolf \_\_\_\_\_ E. \_\_\_\_\_ ENGSTAD  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** Norway

**Residence** Tromso, Norway

**Post Office Address** Strandgata 3, N-9008 Tromso, Norway

**PATENT**

Docket: CU-2652

L 698 184744

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This declaration is of the following type: *(check one applicable item below)*

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- national stage of PCT

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- continuation-in-part (CIP)

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COSMETIC PREPARATIONS

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**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** \_\_\_\_\_ Germany \_\_\_\_\_

**Residence** \_\_\_\_\_ Dusseldorf, Germany \_\_\_\_\_

**Post Office Address** \_\_\_\_\_ Ludolfstr. 13, D-40597 Dusseldorf, Germany \_\_\_\_\_

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**Inventor's signature** \_\_\_\_\_ *Wolf Eisfeld* \_\_\_\_\_

**Date** *09.10.01* \_\_\_\_\_ **Country of Citizenship** \_\_\_\_\_ **Germany** \_\_\_\_\_

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**Post Office Address** Strandgata 3, N-9008 Tromso, Norway \_\_\_\_\_

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Engstad

## **PATENT**

Docket: CU-2652

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### SEND CORRESPONDENCE TO:

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### DIRECT TELEPHONE CALLS TO: (Name and telephone number)

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### Full name of first joint inventor

Ute \_\_\_\_\_ (Given Name) \_\_\_\_\_ (Middle Initial or Name) \_\_\_\_\_ GRIESBACH \_\_\_\_\_ (Family (or Last) Name)

Inventor's signature \_\_\_\_\_

Date \_\_\_\_\_ Country of Citizenship \_\_\_\_\_ Germany \_\_\_\_\_

Residence \_\_\_\_\_ Dusseldorf, Germany \_\_\_\_\_

Post Office Address \_\_\_\_\_ Ludolfstr. 13, D-40597 Dusseldorf, Germany \_\_\_\_\_

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**Full name of second joint inventor**

**Rolf** \_\_\_\_\_ **(Given Name)** \_\_\_\_\_ **(Middle Initial or Name)** \_\_\_\_\_ **WACHTER** \_\_\_\_\_ **(Family (or Last) Name)**

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** Germany

**Residence** Dusseldorf, Germany

**Full name of third joint inventor**

**Achim** \_\_\_\_\_ **ANSMANN** \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature** \_\_\_\_\_

**Date** \_\_\_\_\_ **Country of Citizenship** Germany \_\_\_\_\_

**Residence** Erkrath, Germany

**Full name of fourth joint inventor**

**Bernd** \_\_\_\_\_ **FABRY** \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature**

**Date** **Country of Citizenship** Germany

**Residence** Korschenbroich, Germany

**Full name of fifth joint inventor**

**Wolf** \_\_\_\_\_ **EISFELD** \_\_\_\_\_  
(Given Name) (Middle Initial or Name) (Family (or Last) Name)

**Inventor's signature**

**Date** **Country of Citizenship** Germany

**Residence** Dusseldorf, Germany

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